10

15

20

25

30

PROCESS FOR ACTIVATION OF ALF3 BASED CATALYSTS AND PROCESS FOR ISOMERISING HYDROCHLOROFLUOROCARBONS

The present invention concerns a process for producing an activated AlF₃ catalyst, an activated AlF₃ catalyst and a process for isomerising hydrochlorofluorocarbons.

The invention relates in particular to a process for isomerising 1,1,2-trifluoro-1,2-dichloroethane (CF₂Cl-CHFCl, hereinafter referred to as HCFC-123a or "asymmetrical" isomer) to 1,1,1-trifluoro-2,2-dichloroethane (CF₃-CHCl₂, hereinafter HCFC-123 or "symmetrical" isomer), making use of the activated catalyst. More particularly, it relates to a process for obtaining HCFC-123 with a very low content (less than 0.1%, or 1000 ppm, preferably less than 0.05%, or 500 ppm) of the asymmetrical isomer, HCFC-123a, by isomerisation of the HCFC-123a contained in 123/123a mixtures.

The need for having available an industrial process for preparing HCFC-123 as free as possible from the 123a isomer is well known (see e.g. USP 5,600,037).

However, in the most common industrial process for the preparation of HCFC-123, that is, the fluorination of perchloroethylene with HF in the presence of an appropriate catalyst, the product always contains the 123a isomer, in amounts ranging from some thousands of ppm to some parts %: see ex. WO 95/32168, EP Appl. 609123 (also the 123b isomer CF₂H-CFCl₂ is produced, but in such low amounts that its presence is negligible).

The separation of the isomers with physical methods is exceedingly difficult, due to their similar physico-chemical characteristics (ex. B.P. of $123 = 27.1^{\circ}$ C, of $123a = 28.2^{\circ}$ C).

Several chemical methods have been proposed, to process the 123/123a mixtures and transform the 123a in another compound, more easily separated from 123: for example, disproportionation to HCFC-124 and HCFC-122 (USP 5.414.167), fluorination to HCFC-124 (USP 5.773.671), dehydrochlorination to CTFE (chlorotrifluoroethylene), etc.

Unfortunately, while in each of these methods the 123a isomer is the most reactive of the two, none of them is selective enough to prevent the loss of

-2-

substantial amounts of the symmetrical 123 isomer, which reacts along the 123a, albeit at a slower rate. Moreover, even if 100% selectivity could be achieved, the 123a would be in any case converted to some other compound and is lost.

Several methods have thus been proposed for the catalytic isomerisation of 123a to 123, either in the liquid or in the gas phase.

5

10

15

20

25

30

35

The liquid-phase, homogeneously catalyzed processes are often quite efficient in the isomerisation reaction but suffer several drawbacks, among which the most serious is, as a rule, the low selectivity due to the formation of several undesirable by-products (see e.g. Jap. Appl. 63-85175 of Oct. 16, 1989); a more selective liquid-phase process is disclosed in USP 5,302,766 but it has a very low efficiency: the best result reported is 0.63% residual 123a with a contact time of several hours; more in general, the homogeneous, liquid-phase processes require a more complex and expensive workup of the reaction products than the gas-phase, heterogeneously catalyzed ones.

USP 5,600,037 discloses such a heterogeneous process, in which HCFC-123a is isomerized in the gas phase, on an AlF₃ (aluminium fluoride) solid catalyst. However, in the cited patent the aforementioned goal of less than 0.1% of 123a is never obtained.

Indeed, to obtain a product containing less than 0.1% of 123a, it is necessary to run the isomerisation reaction at a temperature lower than those of the examples of the '037 patent; this is due to the fact that conversion of 123a to 123 is equilibrium-limited and the equilibrium constant between the isomers favours the 123 at low temperatures, the 123a at higher ones. See e.g. USP 5.302.766 (to DuPont), whose results have been confirmed by the data obtained by the Applicant.

Thus, to obtain a product containing a low residual amount of 123a, it is necessary to run the isomerisation reaction at a low temperature; in this case, however, the catalyst deactivates quickly and the useful time on stream is unacceptably short: after a few hours the conversion of 123a to 123 decreases and the residual 123a in the exit stream increases far beyond the desired limit: this is disclosed, e.g., in USP 5,118,887, where in the best mode the conversion of 123a decreases from 99.9 to 85% after a mere three hours on stream; the Applicant's own work confirms these data.

Conversely, at higher temperatures the deactivation is slower but the residual 123a cannot be lower than the equilibrium limit: for example, at 350°C

- 3 -

the deactivation is negligible but the residual 123a cannot be lower than about 0.3%, due to the isomers' equilibrium.

It has now been surprisingly found by the Applicant that these problems can be overcome by using a catalyst based on aluminium trifluoride (AlF₃), which is treated previously to the use in the manner hereinbelow described.

5

10 '

15

20

25

30

35

Consequently the invention concerns a process for producing an activated AlF₃ based catalyst, wherein a crude AlF₃ is treated for more than 5 hours with a gas stream at a temperature from 300°C to 450°C.

When the catalyst obtainable by this process is used, for example, in the gas-phase isomerisation reaction of hydrochlorofluorocarbons such as HCFC—123a, the desactivation of the catalyst is very slow and the reactor needs to be regenerated only at long intervals. The activity of the activated AlF₃ catalyst generally remains at its initial level for at least 10 hours. Often the activity can be preserved for at least 50 hours. In a preferred embodiment, said activity is preserved for at least 100 hours. In a particularly preferred embodiment, said activity is preserved for more than 200 hours.

Excellent results are obtained for the isomerisation of hydrochlorocarbons, in particular of HCFC-123a contained in HCFC-123. Isomeric purity of the product after treatment in the presence of the activated AlF₃ catalyst is generally more than 99.9% mole. Often a purity equal to or greater than 99.95 mole % is achieved.

In the present description, "aluminium trifluoride (AlF₃)" is intended to denote in particular a crystalline solid of such formula, generally obtained from the exhaustive fluorination of aluminium (hydr)oxide (commonly referred to as alumina) with anhydrous hydrogen fluoride (HF), as described e.g. in US 6.187.280 and 6.432.362. During this manufacture of aluminium fluoride, the partially fluorinated aluminas become more and more impervious to further reaction. Consequently, the stoichiometric formula "AlF₃" cannot normally be reached by these methods.

The crude AlF₃ used in the process according to the invention contains generally at least 90 wt. % of stochiometric AlF₃. Preferably, the content of stochiometric AlF₃ is at least 95 wt. %. More preferably the content of stochiometric AlF₃ is at least 96 wt. %. The crude AlF₃ used in the process according to the invention contains generally at most 99.9 wt.% of stochiometric AlF₃. Often, the content of stochiometric AlF₃ is at most 99 wt. %.

The crude AlF₃ used in the process according to the invention has generally a B.E.T. surface area determined by N_2 adsorption equal to or greater than $15 \text{ m}^2/\text{g}$. Often, this specific surface area equal is to or greater than $20 \text{ m}^2/\text{g}$. A specific surface area equal to or greater than $25 \text{ m}^2/\text{g}$ is more particularly preferred. The crude AlF₃ used in the process according to the invention has generally a B.E.T. surface area less than $100 \text{ m}^2/\text{g}$. Often, this specific surface area is equal to or less than $75 \text{ m}^2/\text{g}$. A specific surface area equal to or less than $50 \text{ m}^2/\text{g}$ is more particularly preferred.

5

10

15

20

25

30

35

The crude AlF₃ is preferably chiefly composed by the crystalline phase. The content of crystalline phase as determined by X-ray diffraction and comparison of the relative peak intensity is generally at least 60%. Preferably, this content is at least 70%. The content of crystalline phase as determined by X-ray diffraction and comparison of the relative peak intensity is generally less than 100%. Preferably, this content is at most 85%.

When the crude AlF₃ is obtained by fluorination of alumina, the alumina used as a starting material is preferably in the form of the hydrated aluminium oxide known as boehmite and may optionally contain a minor component of silicon oxide (silica). Such aluminas or silico-aluminas are commercial products, for example the Pural® and Siral® grades of the Sasol (ex Condea) firm.

Moreover, if the catalyst is to be used in a fluid-bed reactor, the alumina has a particle size distribution compatible with this use, as it is well known by the expert in the art.

Beyond the cited patents, both ΛIF_3 as such and the method for its preparation are well known to the art, see e.g. FR 1.383.927.

In the process according to the invention, the temperature of the treatment is preferably equal to or greater than 330°C. A temperature equal to or greater than 350°C is more particularly preferred. In the process according to the invention, the temperature of the treatment is preferably equal to or less than 420°C. A temperature equal to or less than 400°C is more particularly preferred.

In the process according to the invention, the duration of the treatment is preferably equal to or greater than 6 hours. A duration equal to or greater than 10 hours is more particularly preferred. In the process according to the invention, the duration of the treatment is generally equal to or less than 50 hours. A duration equal to or less than 20 hours is more particularly preferred.

In the process according to the invention, the pressure of the treatment is generally equal to or greater than about 1 bar. Atmospheric pressure (about 1 bar) is preferred. In the process according to the invention, the pressure of the treatment is generally equal to or less than about 10 bar. A pressure equal to or less than about 5 bar is preferred.

5

10

15

20

25

30

35

In the process according to the invention, the contact time of the treatment defined as the ratio between the volume of the reactor and the flow of treatment gas is generally equal to or greater than about 10s. A contact time equal to or greater than about 30s is preferred. In the process according to the invention, the contact time is generally equal to or less than about 100s. A contact time equal to or less than about 50s is preferred.

In the process according to the invention, the gas stream contains often at least one gas selected from air, oxygen, hydrogen fluoride, halogenated hydrocarbon or inert gases. Among the inert gases, particular mention may be made of nitrogen and noble gases such as helium and argon. Among the halogenated hydrocarbons, particular mention may be made of chlorohydrocarbons, fluorohydrocarbons and hydrochlorofluorocarbons. Among the latter, 1,1,1-trifluoro-2,2-dichloroethane and its mixtures with the isomer 1,1,2-trifluoro-1,2-dichloroethane is particularly preferred.

If a halogenated hydrocarbon stream is used, it can consist essentially of the halogenated hydrocarbon or it can optionally be diluted, for example with an inert gas.

If the activated AlF₃ catalyst is intended for use for the isomerisation of a hydrochlorofluorocarbon, the activation comprises, in a preferred embodiment, an activation treatment with the hydrochlorofluorocarbon mixture to be isomerised.

In a particularly advantageous embodiment of the process according to the invention, the treatment with the gas stream comprises at least 2 treatment steps with different gases, preferably selected from those mentioned above.

In this embodiment, the treatment with the gas stream comprises preferably at least a treatment step carried out with an anhydrous hydrogen fluoride stream. In this case, the duration of the treatment with the anhydrous hydrogen fluoride stream is generally equal to or greater than 4 hours. Preferably the duration of this treatment is equal to or greater than 6 hours. In this case, the duration of the treatment with the anhydrous hydrogen fluoride

10

15

20

25

30

35

stream is generally equal to or less than 16 hours. Preferably the duration of this treatment is equal to or less than 12 hours.

In a particularly preferred embodiment, the temperature of the treatment with anhydrous hydrogen fluoride is about $350^{\circ}\text{C} \pm 25^{\circ}\text{C}$ or $360^{\circ}\text{C} \pm 25^{\circ}\text{C}$.

In a first particularly preferred mode of carrying out the said embodiment, the treatment with the gas stream comprises

- (a) a treatment with an inert gas stream for at least 2 hours
- (b) optionally, a treatment with an anhydrous hydrogen fluoride stream, in particular as described herebefore,
- (c) a treatment with a hydrochlorofluorocarbon containing stream for more than 3 hours.

In this mode, the duration of treatment step (a) is preferably at least 4 hours. The duration of treatment step (a) is generally at most 12 hours. The duration of treatment step (a) is preferably at most 8 hours.

In this mode, the duration of treatment step (c) is preferably at least 7 hours. The duration of treatment step (c) is generally at most 50 hours. The duration of treatment step (c) is preferably at most 25 hours. It should be understood that in this mode the exit gases can be monitored via GC, and the treatment is deemed complete when a constant or declining conversion of hydrochlorofluorocarbon is attained. The time necessary to complete this part of the activation is considerably shortened, if it is preceded by the HF treatment. In a particularly preferred embodiment, the temperature in treatment step (c) is about $350^{\circ}\text{C} \pm 25^{\circ}\text{C}$.

In a second particularly preferred mode of carrying out the said embodiment, the treatment with the gas stream comprises

- (a) a treatment with an air stream for at least 2 hours
- (b) a treatment with an anhydrous hydrogen fluoride stream for at least 4 hours.

In this mode, the duration of treatment step (a) is preferably at least 4 hours. The duration of treatment step (a) is generally at most 12 hours. The duration of treatment step (a) is preferably at most 8 hours.

In a particularly preferred embodiment of this mode, the temperature in treatment step (a) is about $390^{\circ}\text{C} \pm 25^{\circ}\text{C}$.

In this mode, the duration of treatment step (b) with anhydrous hydrogen fluoride is as described above.

It is understood that the information of the general description of the process applies also to each of the particular embodiments described herebefore.

The invention concerns also an activated AlF₃ catalyst, obtainable according to the process according to the invention.

5

10

15

20

25

30

35

The activated catalyst obtained according to the process according to the invention may be either unloaded and stored, or immediately used in the same reactor e.g. for the isomerisation of hydrochlorofluorocarbons such as HCFC-123a.

In this case the temperature is suitably adjusted to the value required to obtain the desired content of isomer.

The invention concerns also a process for the isomerisation of a hydrochlorofluorocarbon, wherein the hydrochlorofluorocarbon is contacted with the activated catalyst according to the invention.

In the isomerisation process according to the invention, the hydrochlorofluorocarbon is preferably in the vapor state.

The isomerisation process according to the invention can be carried out in a fixed bed reactor or in a fluidized bed reactor. A fluidized bed reactor is preferred.

Hydrofluorocarbons which can suitably be isomerised by the isomerisation process according to the invention include, amongst others, 1,1,2-trifluoro-1,2,-dichloroethane, 1,1,2-trifluoro-1-chloroethane, 1,2-difluoro-1-chloroethane, 1, fluoro-1,2,-dichloroethane, 1,1,3-trifluoro-1,3,-dichloropropane and 1,1,3,3-tetrafluoro-3-chloropropane. The isomerisation of 1,1,2-trifluoro-1,2,-dichloroethane is particularly preferred. The following description is specifically drawn to the isomerisation reaction of 1,1,2-trifluoro-1,2,-dichloroethane, it should however be understood that the subject or similar conditions apply also to other hydrochlorofluorocarbons.

In a preferred embodiment of the isomerisation process according to the invention, the hydrochlorofluorocarbon introduced into the reactor comprises a mixture of 1,1,1-trifluoro-2,2,-dichloroethane (123) and 1,1,2-trifluoro-1,2,-dichloroethane (123a).

In this embodiment, the 123/123a mixture fed to the reactor contains generally at least 0.3 wt. % of 123a with reference to the total weight of the 123/123a mixture. Often this content is equal to or greater than about 0.5 wt. %. More frequently, this content is equal to or greater than about 1 wt. %. In

-8-

this embodiment, the 123/123a mixture fed to the reactor contains generally at most 50 wt. % of 123a with reference to the total weight of the 123/123a mixture. Often this content is equal to or less than about 20 wt. %. More frequently this content is equal to or less than about 15 wt. %.

5

10

15

20

25

30

35

In this embodiment, the organic feed introduced into the isomerisation reaction may contain, beside the 123 isomers, also substantial amounts of different compounds, such as those found as by-products in a 123 production process such as contemplated herebefore. It has been found that these compounds do not adversely affect the isomerisation process according to the invention. The organic feed introduced into the isomerisation reaction may typically contain up to approx. 20% by weight of by-products and 80% weight of 123 isomers.

Surprisingly, no appreciable difference in the results of the isomerisation reaction has been found, whichever the isomeric composition of the organic feed. The isomerisation process according to the invention is flexible enough to treat, for example, all possible isomeric compositions of an HCFC-123 produced e.g. via fluorination of perchloroethylene, under industrially relevant conditions. If a sufficient conversion of 123a to 123 is not reached because of a very high content of 123a in the feed, this can be overcome by adjusting the recycle loop of the reactor until the appropriate inlet composition has been achieved.

In the isomerisation process according to the invention, the organic feed may optionally be diluted in an inert gas, such as described above. If present, the inert gas fraction of the total stream is generally at least 5% by volume. In this case, this fraction is preferably at least 10% by volume. If present, the inert gas fraction of the total stream is generally at most 90% by volume. In this case, this fraction is preferably at most 50% by volume.

In the isomerisation process according to the invention, the temperature is preferably equal to or greater than 150°C. A temperature equal to or greater than 180°C is more particularly preferred. A temperature about equal to or greater than 200°C is most particularly preferred. In the isomerisation process according to the invention, the temperature is preferably equal to or less than 260°C. A temperature equal to or less than 230°C is more particularly preferred. A temperature about equal to or less than 220°C is most particularly preferred.

The pressure of the reactor does not adversely affect the isomerisation reaction, provided it is not so high to cause the condensation of the hydrochlorofluorocarbon at the reaction temperature.

In the isomerisation process according to the invention, the pressure is generally equal to or greater than about 1 bar. A pressure equal to or greater than about 2 bar is preferred. In the isomerisation process according to the invention, the pressure is generally equal to or less than about 10 bar. A pressure equal to or less than about 5 bar is preferred.

While the isomerisation of 123a is not adversely affected by the pressure of the reactor, it has been found, surprisingly, that a moderately high pressure has a beneficial effect on the formation of unsaturated by-products. In particular, when the process is run at a pressure which is moderately higher than the atmospheric, preferably at 2 to 5 bar abs., the formation of the unsaturated by-product CFC-1112a (1,1-dichloro-2,2-difluoroethylene, CF₂=CCl₂) is greatly reduced or even eliminated altogether. This is desirable since CFC-1112a is both toxic and difficult to separate from HCFC-123.

The invention concerns in consequence also a method for the isomerisation of 1,1,2-trifluoro-1,2,-dichloroethane wherein the 1,1,2-trifluoro-1,2,-dichloroethane, preferably in the vapor state, is contacted with an isomerisation catalyst under a pressure of from 2 to 5 bar.

The particular embodiments of the method according to the invention correspond to those of the isomerisation process according to the invention.

The instant invention is illustrated in a non-limitative manner by the following examples:

25 Example 1 - Activation of the catalyst

5

10

15

20

30

A sample of 9.0 g of aluminium trifluoride in form of a fine powder obtained by fluorination of a SIRAL® 1.5 silico-alumina using the general fluorination procedure described in US 6.187.280 (examples) was loaded into a tubular reactor of 3/8 inches o.d. (ca 15 mm internal diameter), heated by means of an electric oven, and brought to the temperature of 350°C in an helium flow.

After 4 hours at 350°C, a flow of 2.1 scc (cubic centmeters at 20°C and atmospheric pressure)/min (0.8 g/h) of HCFC-123, containing 11.9% 123a, was added to the feed. The pressure in the reactor was atmospheric, the total flow of He and organics was 7 scc/min.

20

25

The gas exiting from the reactor was sampled an analysed via GLC; the flow was monitored at about 40min intervals, witnessing an increase of the conversion of 123, which reached a maximum after ca. 9 hrs on stream.

Both isomers 123 were converted to several disproportionation products: the following analysis, at 9 hrs on stream, is typical of the composition of the products when the maximum conversion had been attained:

	HCFC-124*	9.7% weight
	HCFC-133a	1.7
	CFC-114*	3.6
10	CFC-113*	0.6
	HCFC-123a	0.3
	HCFC-123	76.8
	CFC-1112*	0.1
	HCFC-1111*	5.3
15	HCFC-122*	1.0
	others	0.9

^{*}Mixture of isomers

The residual 123a/123 content was stable at ca. 0.35%, which is the about equilibrium limit at 350°C.

When the conversion of 123 had reached the maximum, the temperature of the reactor is lowered to the desired isomerisation temperature.

Example 2 - Isomerisation of 123a at 220°C (fixed bed reactor)

Continuing from ex. 1, the temperature of the reactor was lowered to 220°C and the sampling was resumed. The following analysis of the obtained products shows that at this temperature the formations of by-products was negligible and the residual 123a/123 content was down to ca. 500 ppm

	HCFC-123a	0.05
	HCFC-123	99.1
	CFC-1112a	n.d.
30	others	0.85

The selectivity (measured as the fraction of 123 recovered) was equal to 99.2%; the above results were obtained consistently during the whole run (about 6 hours), with no deactivation of the catalyst.

15

25

30

Example 3 - Isomerisation of 123a at 200°C

After an interval, the testing was resumed with the same catalyst used in Examples. 1 and 2, which was briefly reactivated with a flow of He/123 at 350°C, then the temperature of the reactor was lowered to 200°C. The same stream of ex. 2 was fed to the reactor at atmospheric pressure.

The products were sampled during a 4 hrs period. Their composition was substantially stable throughout the run, with the following analytical result:

	HCFC-123a	0.04
	HCFC-123	99.5
10	CFC-1112a	n.d.
	others	0.45

The residual 123a/123 was about 400 ppm, lower than in ex. 2, and the selectivity as well was increased, to about 99.5%.

Example 4 - Isomerisation of 123a at 200°C with an HCFC-123a containing a lower amount of HCFC-123a

Example 3 was reproduced (with a fresh catalyst sample, previously activated as in Ex. 1) with the only change that the HCFC-123 mix contained 3.8% wt. of the asymmetrical isomer; no meaningful difference in the results was detected, as results from the following analysis of the products:

20	HCFC-123a	0.04
	HCFC-123	99.6
	CFC-1112a	n.d.
	others	0.035

The residual 123a content in the products was extremely low and the overall selectivity of the reaction was the same as in Ex. 3.

Example 5 - Comparison with Example 4

A fresh sample of the same AlF₃ used in Ex. 1-4 has been used in comparison test: thus, 10 g of catalyst were loaded in the same reactor and dehydrated at 350°C in an He flow for only 1 hr. then activated with an HCFC-123 flow at 350°C in the same conditions as in Ex. 4 but for two hours only.

The temperature was then lowered to 210°C and the same He/123 flow of Ex. 4 (with 3.8% 123a) was fed to the reactor.

The conversion of 123a decreased fast and after 5 hrs on stream the content in the product gas had reached 0.26%, far higher than the desired value.

10

15

20

25

30

35

Thus omitting the activation according to the invention resulted also in an unacceptably short useful life of the catalyst.

Example 6 - Fixed bed reaction with pelletized catalyst

In this example a pelletized catalyst was employed. This catalyst is especially suitable for the use in industrial-scale fixed-bed reactors.

400 g (375 cc) of AlF₃ pellets were loaded in a tubular reactor, 50 mm i.d., equipped with a fritted bottom and thermally insulated. In this setup, the reactor was operated in so-called "adiabatic" mode, where the heat is furnished to the reaction feed in an exchanger before entering the reactor.

The pressure in the reactor was atmospheric both during the activation and the isomerisation test.

The reactor was brought to 360°C in a nitrogen flow, then the catalyst was first treated in an HF flow, purged from HF again in a nitrogen flow and finally activated with a 123 flow for a period of 20 hrs, in the following conditions:

Temperature 350°C, 123 feed 0.266 mol/h (41 g/h) diluted with 21 sL/h of nitrogen; the nominal contact time at 350°C and atm. pressure was 33 s; the HCFC-123 contained 0.3% (3000 ppm) of the asymmetric isomer.

The reactor temperature was then adjusted to 240°C while the same flow was fed to the reactor, thus realizing a contact time of 40 s.

The reaction products were sampled and analyzed and a residual 123a content of 0.025 (250 ppm) was found; this result remained constant for the whole duration of the test, 30 h on stream.

Example 7 - Isomerisation of 123a in a fluid-bed reactor

400 g (380 cc) of fresh AlF₃, with a particle size distribution suitable for the use as a fluid-bed catalyst, were loaded in a tubular reactor of 50 mm equipped with a fritted bottom and heated by means of electric resistors. The catalyst was heated up to 390°C in a nitrogen flow, then treated for 4 hr with an air flow at 390°C. Then the temperature of the reactor was set to 360°C, the air flow was discontinued and replaced by an HF flow. After 10 hrs the HF flow was stopped and the reactor was purged in nitrogen flow for a few hrs, while the temperature was gradually lowered to 240°C.

When the reactor temperature had stabilized at 240°C, a reaction mixture of the following composition was fed to the reactor: 40.7 g/h (0.266 mol/h) of HCFC-123 (containing 0.3% HCFC-123a) and ca. 15 NL/h (0.663 mol/h) of nitrogen.

15

35

The nominal contact time was 35 s at the reaction temperature and atmospheric pressure.

The products were sampled and analyzed via GC with the following result:

5	HCFC-124+124a	1.61	mol %
	HCFC-123a	0.067	
٠.	HCFC-123	97.47	
	CFC-1112a	0.33	
	others	0.52	

The catalytic activity remained stable during the whole run (20 hrs), with no sign of deactivation. The residual 123a content remained on average at about 700 ppm and the selectivity around 97.5%.

Example 8

The test of ex. 7 was continued at a lower temperature, without any other alteration. The temperature of the reactor was set at 210°C without even stopping the reactant flow. The flow was not adjusted to the new temperature, thus the contact time became slightly higher than in. ex. 7.

The residual 123a dropped to 400 ppm and the conversion of 124 and formation of 1112 dropped too:

20	HCFC-124+124a	0.62 r	nol %
	HCFC-123a	0.034	
	HCFC-123	98.99	
	CFC-1112a	0.14	
	others	0.22	

Both the selectivity (about 99%) and the residual content of 123a isomer about 350-400 ppm remained substantially constant for the duration of the test, 70 hrs on stream.

Example 9

Example 7 was repeated with a fresh catalyst sample, which was activated in the same manner; at a temperature of 240°C, a flow of HCFC-123 was fed to the reactor under identical conditions as in example 7, excepted that the 123a content in the feed stream was 1.4% wt. The residual 123a in HCFC-123 was stable at about 700 ppm.

HCFC-124+124a	2,23	mol %
HCFC-123a	0.063	

15

20

-		
-	14	-

HCFC-123	96.26
CFC-1112a	0.55
others	0.90

Example 10

In an analogous manner as described in examples 7 and 8, the reactor temperature was lowered to 210°C at the end of the procedure of example 9 without stopping the reactant flow. The residual 123a dropped to about 400 ppm and the content of by-products as well.

	HCFC-124+124a	0.57	mol %
10	HCFC-123a	0.038	
	HCFC-123	99.14	
* -	CFC-1112a	0.10	
	others	0.15	

The catalyst showed no deactivation for the whole duration of Example 10, ca 35 hrs on stream

Example 11 - Isomerisation of 123a at superatmospheric pressure in a fluid-bed reactor

In the same experimental setup of example 7, 400 g of fresh AlF₃ was loaded and activated in the same manner, as in that example.

After the activation, the temperature was set at 240°C and the reactor pressure at 3 atm. absolute (2 atm. above atmospheric), and a mixture of 40.7 g/h (0.266 mol/h) of HCFC-123 (containing 0.3% 123a) and ca. 63 NL/h of nitrogen, was supplied to the reactor.

The following results were obtained:

25	HCFC-124	1.37	mol %
	HCFC-123a	0.062	
	HCFC-123	98.42	
	CFC-1112a	0.20	
	others	0.05	

These results remained constant for 160 hrs on stream, during which the catalyst showed no sign of deactivation.

It is readily apparent that the pressure increase had no adverse effect on the isomerisation of 123a, while it is quite effective in reducing the byproduction of 1112a, which is a toxic and undesirable compound.

Example 12

The test of example 1 was continued without regeneration of the catalyst; the total feed streams remained the same but the 123a content in HCFC-123 was increased to 1.5% wt. with the following result:

5	HCFC-124	0.68	mol %
	HCFC-123a	0.055	
	HCFC-123	98.90	
	CFC-1112a	0.25	
	others	0.12	

This example shows that when using a 123 feed with an higher content of 123a, the results were not affected and the residual 123a in the products reached still the equilibrium limit. These results remained constant up to the end of the test i.e. more than 200 hrs on stream from the beginning of example 1. A decrease in time of the side reaction to HCFC-124 has also been observed.

15 Example 13

20

The catalyst used in Examples 1 and 12 was regenerated by means of an air treatment at 390°C for 4 hours followed by fluorination with HF at 360°C for 12 hours.

After the regeneration the 123/123a (ca. 4000 ppm 123a) flow was restored with the same pressure and feedrates as in the examples 1 and 12, while the reaction temperature was set at 220°C. At this lower temperature, the residual 123a in 123 decreased to 400 ppm average and remained stable for the whole duration of the test, 140 hrs. The 1112a and other byproducts decreased as well:

25	HCFC-124	0.58 mol %
	HCFC-123a	0.035
	HCFC-123	99.29
	CFC-1112a	0.06
	others	0.04

30 Example 14

The catalyst used in examples 11-13 was regenerated as above described and the pressure in the reactor was set to 5 ata (4 bar gauge). The temperature was 220°C.

- 16 -

To conserve the same volumetric contact time, both the 123 and nitrogen feedrates were increased to:

$$123 = 0.443 \text{ mol/h} = 68 \text{ g/h}$$

nitrogen = 104 sL(liter at 20°C and atmospheric pressure)/h

5 The following result was obtained:

HCFC-124 0.05 HCFC-123a 0.06 HCFC-123 99.89

Any other byproduct was lower than the analytical threshold. Thus the pressure is shown to have a very beneficial effect indeed.

20

CLAIMS

- 1 Process for producing an activated AlF₃ based catalyst, wherein a crude AlF₃ is treated for more than 5 hours with a gas stream at a temperature from 300°C to 450°C.
- 5 2 Process according to claim 1, wherein the temperature is from 350°C to 400°C.
 - 3 Process according to claim 1 or 2, wherein the crude AlF₃ is treated with the gas stream for from 6 to 50 hours.
- 4 Process according to any one of claims 1 to 3, wherein the crude AlF₃, contains at least 95 wt.% of stochiometric AlF₃.
 - 5 Process according to any one of claims 1 to 4, wherein the crude AIF₃ has a B.E.T surface of at least 25 m²/g.
 - 6 Process according to any one of claims 1 to 5, wherein the gas stream contains at least one of air, hydrogen fluoride, halogenated hydrocarbon or inert gas.
 - 7 Process according to any one of claims 1 to 6, wherein the treatment with the gas stream comprises at least 2 treatment steps with different gases.
 - 8 Process according to claim 7, wherein the treatment with the gas stream comprises
 - (a) a treatment with an inert gas stream for at least 4 hours
 - (b) optionally, a treatment with an anhydrous hydrogen fluoride stream
 - (c) a treatment with a hydrochlorofluorocarbon-containing stream for more than 1 hour.
- 9 Process according to claim 7, wherein the treatment with the gas stream comprises
 - (a) a treatment with an air stream for at least 2 hours
 - (b) a treatment with an anhydrous hydrogen fluoride stream for at least 4 hours.

- 10 Activated AlF₃ catalyst, obtainable according to the process of any one of claims 1 to 9.
- 11 Process for the isomerisation of a hydrochlorofluorocarbon, wherein the hydrochlorofluorocarbon is contacted with the catalyst according to claim 10.
- 5 12 Process according to claim 11 wherein the hydrochlorofluorocarbon is in the vapor state.
 - 13 Process according to claim 11 or 12, wherein the hydrochlorofluorocarbon comprises a mixture of 1,1,1-trifluoro-2,2-dichloroethane and 1,1,2-trifluoro-1,2-dichloroethane.
- 10 14 Process according to claim 13 wherein the isomerisation is carried out at a temperature of 180 to 220°C.
 - 15 Method for the isomerisation of 1,1,2-trifluoro-1,2-dichloroethane wherein the 1,1,2-trifluoro-1,2-dichloroethane, preferably in the vapor state, is contacted with an isomerisation catalyst under a pressure of from 2 to 5 bar.

International Application No
T/EP2004/051624

A. CLASSIFICATION OF SUBJECT MATTER IPC 7 C07C17/358

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols) IPC 7 C07C

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

EPO-Internal

Category °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
x	EP 0 881 201 A (AUSIMONT SPA)	1-4,6-8,
	2 December 1998 (1998-12-02) page 2, line 53 - line 58	10
].	claims 1,4; examples 1A,1B	and the second s
χ	LIC 2 707 221 A (CACTELLANDALET ALL)	1 2 6 0
^	US 3 787 331 A (CASTELLAN A ET AL) 22 January 1974 (1974-01-22)	·· 1-3;6-8; 10-12
	column 4, line 39 - line 49; examples 2,5;	10 12
1	table 4	•
x	US 5 731 481 A (LANTZ ANDRE ET AL)	1-8,10
	24 March 1998 (1998-03-24)	
	examples 1A,1B,2A,2B	
x	US 5 616 820 A (CHEMINAL BERNARD ET AL)	1-3,6-8,
	1 April 1997 (1997-04-01)	10-13
	column 6, line 8 - line 35; examples A,B; table 1	
	_/ _	

Further documents are listed in the continuation of box C.	Patent family members are listed in annex.				
Special calegories of cited documents: A* document defining the general state of the art which is not considered to be of particular relevance 'E* earlier document but published on or after the international filing date 'L* document which may throw doubts on priority claim(s) or	 "T* later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art. "&" document member of the same patent family 				
which is cited to establish the publication date of another citation or other special reason (as specified) "O" document referring to an oral disclosure, use, exhibition or other means "P" document published prior to the international filing date but later than the priority date claimed					
Date of the actual completion of the international search 17 September 2004	Date of mailing of the international search report 28/09/2004				
Name and mailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Authorized officer Holzwarth, A				

International Application No
T/EP2004/051624

	ntion) DOCUMENTS CONSIDERED TO BE RELEVANT	
Category °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	US 5 600 037 A (CUZZATO PAOLO ET AL) 4 February 1997 (1997-02-04) cited in the application example 3; table 3	1-3,7, 9-13
	· · · · · · · · · · · · · · · · · · ·	
	•	
į		
		·
	·	

Information on patent family members

International Application No
T/EP2004/051624

							4/ 031024
i i	atent document d in search report		Publication date		Patent family member(s)		Publication date
EP	0881201	A	02-12-1998	IT DE DE EP JP US	MI971237 69807980 69807980 0881201 10330298 6300529	D1 T2 A1 A	27-11-1998 24-10-2002 31-07-2003 02-12-1998 15-12-1998 09-10-2001
US	3787331	A	22-01-1974	DK AT BE CH DE FR GB IL JP NO SE YU ZA	138673 292641 740090 525710 1950804 372331 2020296 1283386 33144 55012297 6914962 125479 361135 254669	B A A A1 A5 A A B B B B B	16-10-1978 10-09-1971 10-04-1970 31-07-1972 29-10-1970 16-02-1972 10-07-1970 26-07-1972 29-06-1973 01-04-1980 14-04-1970 18-09-1972 22-10-1973 28-02-1975 27-01-1971
US	5731481	Α	24-03-1998	FR AU AU CA DE DE ES JP JP KR NO PT DK GR IE MX	2669022 115532 641292 8779991 2055281 69105924 69105924 0486333 2066394 915341 2053012 4288027 7080796 9506521 914107 99489 486333 3015222 913932 9102049	T T B2 B2 A1 B1 T2 B A1 B A1 B A B B1 B1	15-05-1992 15-12-1994 16-09-1993 14-05-1992 14-05-1992 26-01-1995 27-07-1995 20-05-1992 01-03-1995 14-05-1992 10-05-1996 13-10-1992 30-08-1995 14-05-1995 30-09-1992 01-05-1995 31-05-1995 20-05-1995 20-05-1992 08-07-1992
US	5 5616820	A	01-04-1997	FR AU BR CA CN DE DE EP GR JP KR	2700770 663816 5395194 9400338 2113511 1101337 69400342 69400342 0609124 2091097 3021203 2510126 6247883 9700398	5 B2 4 A 8 A 1 A1 7 A ,B 2 D1 2 T2 4 A1 7 T3 5 B2 3 A	29-07-1994 19-10-1995 18-08-1994 16-08-1994 28-07-1994 12-04-1995 05-09-1996 23-01-1997 03-08-1994 16-10-1996 31-12-1996 26-06-1996 06-09-1994 09-01-1997

Information on patent family members

International Application No T/EP2004/051624

Patent document cited in search report	Publication date	Patent family member(s)		Publication date
US 5600037 A	04-02-1997	IT	1251957 B	27-05-1995
		ΑT	151403 T	15-04-1997
		ΑU	652080 B2	11-08-1994
1		ΑU	2703892 A	22-04-1993
		BR	9203978 A	27-04-1993
		CA	2080894 A1	19-04-1993
		CZ	9203110 A3	16-06-1993
		DE	69218870 D1	15-05-1997
		DE	69218870 T2	06-11-1997
		EP	0537759 A2	21-04-1993
		ES	2103021 T3	16-08-1997
		GR	3023176 T3	30-07-1997
		JP	3231431 B2	19-11-2001
		JP	6128180 A	10-05-1994
		KR	236863 B1	15-01-2000
		RU	2089534 C1	10-09-1997
		ZA	9207873 A	27-04-1993